The Functional Organization of the Ventral Visual Pathway in Humans

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1. Introduction

We can recognize an object within a fraction of a second, even if we have never seen that particular object before, and even if we have no advance information about what kind of object it might be (Potter, 1976; Thorpe, Fize & Marlot, 1996). The cognitive and neural mechanisms underlying this remarkable ability are not well understood, and current computer vision algorithms still lag far behind human performance. One promising strategy for attempting to understand human visual recognition is to characterize the neural system that accomplishes it: the ventral visual pathway (VVP), which extends from the occipital lobe into inferior and lateral regions of the temporal lobe. Here we describe key results from the last fifteen years of neuroimaging research on humans that have begun to elucidate the general organization and functional properties of the cortical regions involved in visually perceiving people, places, and things.

The central finding from this now substantial body of work is that the VVP is not homogeneous, but instead a highly differentiated structure containing a set of regions each with its own distinct functional profile. These regions include the fusiform face area (FFA), which responds selectively to faces (Kanwisher, McDermott & Chun, 1997; McCarthy, Puce, Gore & Allison, 1997), the parahippocampal place area (PPA), which responds selectively to places (Epstein & Kanwisher, 1998), the extrastriate body area (EBA), which responds selectively to bodies (Downing, Jiang, Shuman & Kanwisher, 2001), the lateral occipital complex (LOC), which responds to object shape (Kanwisher, Woods, Iacoboni & Mazziotta, 1997; Malach et al., 1995) largely independent of object category, and the visual word form area (VWFA), which responds selectively to both visually presented words and consonant strings (Baker, Hutchison & Kanwisher, 2007; Cohen et al., 2000). Each of these regions is present in approximately the same location in virtually every healthy subject. These regions, and their cohorts (e.g., the occipital face area, OFA; see Section 4), constitute the fundamental machinery of high-level visual recognition in humans. An understanding of the function of each of these regions is likely to provide important clues about how visual recognition works.

In this chapter, we first describe the best-established functionally specific regions in the VVP, and then the ongoing controversies about the degree of such specialization. We then review the available evidence on the functional organization of the entire VVP, and finally we consider the computational advantages that may be afforded by having specialized regions in the first place.

2. The best-established functionally specific regions in the ventral visual pathway

Figure 1 shows the main functionally distinct regions of the VVP. We begin by focusing on the FFA, PPA, EBA, and LOC, because these are the best-established regions in this pathway, in several respects. First, each of these regions has been found consistently in a large number of studies and labs; although their theoretical significance can be debated, their existence cannot. Second, the category selectivity by which each region is defined is
not merely statistically significant, but also large in effect size: Each of these regions responds about twice as strongly to any stimulus from its preferred category as to any nonpreferred stimulus category. Effect size is often ignored in the brain imaging literature, but it should not be, as it determines the strength of the inference you can draw: If you know how to double the response of a region, you generally have a better handle on its function than if you merely know how to change its response by a small amount. Third, the fact that these regions can be found easily and now algorithmically (Julian, Fedorenko, Webster & Kanwisher, 2012) in any healthy subject has made possible a “region of interest” (ROI) research strategy whereby the region is first functionally identified in each subject individually in a short “localizer” scan, and then characterized in rich detail in subsequent experiments, including not just its response profile but the information it represents.

The approach to neuroimaging in which cognitive functions are assigned to brain regions has been widely maligned as “mere phrenology”, as if the label itself is an argument against the whole enterprise. But name-calling is not argumentation, any more in science than in the schoolyard. There is nothing fundamentally wrongheaded with the effort to characterize the cognitive functions of particular brain regions— it is an empirical question how successful this research program will turn out to be. The problem is rather that most neuroimaging studies neither robustly establish the existence of a functionally specific region, nor precisely identify its function. When a brain region is identified based on just a few conditions in a single study, the evidence will necessarily be weak and the functional characterization of the region will be sparse and unsatisfying. But when the same brain region has been identified in many labs, and when each of those labs has measured the functional response of that region in numerous conditions, each testing a different hypothesis about the representations it contains, then we can begin to approach the kind of rich cognitive (and perhaps some day computational) characterization of that brain region that will be of real theoretical significance to cognitive science. Thus, brain imaging can discover not just the locations of cognitive functions (who cares?), but more fundamentally it can discover and characterize those cognitive functions themselves. That is, brain imaging can exploit the functional specificity of the brain to discover the cognitive architecture of the mind. Although still very much a work in progress, this enterprise is most advanced for the regions we begin with here: the FFA, PPA, EBA, and LOC.

a. The Fusiform Face Area (FFA). The FFA is the region of the mid-fusiform gyrus (on the bottom surface of the cerebral cortex, just above the cerebellum) that responds significantly more strongly when subjects view faces than when they view objects (Kanwisher et al., 1997; McCarthy et al., 1997; Puce, Allison, Asgari, Gore & McCarthy, 1996). The precise shape of the FFA varies across subjects, consisting of two regions in some subjects (Weiner & Grill-Spector, 2012), but is present in nearly all subjects and highly reproducible within each (Peelen & Downing, 2005). Earlier neuroimaging work (Haxby et al., 1994; Puce, Allison, Gore & McCarthy, 1995) had shown activations in this general region for faces (compared to scrambled faces, textures, and letterstrings), but those contrasts left open the possibility that this region was engaged in processing generic object shape. The subsequent evidence for the specificity of this region for faces
per se came from the discovery that this region responds similarly and strongly to a wide variety of face images (Kanwisher & Yovel, 2006), including photos of familiar and unfamiliar faces, schematic faces, cartoon faces, and cat faces (Kanwisher & Barton, 2010), as well as faces presented in different sizes, locations, and viewpoints (Axelrod & Yovel, 2012; Grill-Spector et al., 1999; Schwarzlose, Swisher, Dang & Kanwisher, 2008), and much less strongly to nonface stimuli. Extensive evidence now rejects alternative hypotheses proposed earlier that the FFA is more generally engaged in fine-grained discrimination of exemplars of any category, or of any category for which the subject has gained substantial expertise (McKone & Robbins, 2010; Tarr & Gauthier, 2000; Yovel & Kanwisher, 2004). Consistent with the evidence from fMRI, face-selective responses have been observed in approximately the same location in subdural electrode recordings from the brains of subjects undergoing presurgical mapping for epilepsy treatment (Allison, Puce, Spencer & McCarthy, 1999; McCarthy, Puce, Belger & Allison, 1999; Puce, Allison & McCarthy, 1999), and lesions in approximately this location can produce selective deficits in face perception (Kanwisher & Barton, 2010). Thus, as discussed further in Section 4, the FFA appears to be quite selectively engaged during the perception of faces.

Answering the question of what exactly the FFA does with faces has been more difficult. Importantly, the magnitude of the FFA response is correlated trial-by-trial with success both in detecting the presence of faces, and in identifying individual faces (Grill-Spector & Malach, 2004), but not with detection or identification of nonface objects, and the extent of face selectivity in the fusiform is correlated with behavioral ability at face (but not object) identification across subjects (Furl, Garrido, Dolan, Driver & Duchaine, 2011). In terms of the kind of information represented in the FFA, current evidence indicates that the FFA is sensitive to multiple aspects of face stimuli including face parts (eyes, noses, and mouths), the T-shaped configuration of those features, and external features of faces (Liu, Harris & Kanwisher, 2010). Further, FFA responses show some invariance across changes in stimulus position and image size (Schwarzlose et al., 2008; but see Yue, Cassidy, Devaney, Holt & Tootell, 2011). Using multivoxel pattern analysis (MVPA), a recent study (Axelrod & Yovel, 2012) found similar neural representations for mirror-symmetric views of faces, but not for other changes in viewpoint, while other studies using fMRI adaptation have found a moderate invariance for viewpoint changes (up to 30°) for familiar (celebrity) faces (e.g., Ewbank & Andrews, 2008). Finally, the FFA exhibits neural correlates of well-established behavioral signatures of face perception (McKone & Robbins, 2010), including sensitivity to differences in face identity for upright but not inverted faces (Yovel & Kanwisher, 2005) and sensitivity to holistic information in upright but not inverted faces (Schiltz & Rossion, 2006). Thus, the FFA appears to represent perceptual information about face shape in a fashion partially invariant to image changes, and to reflect the well-known behavioral signatures of face-specific processing.

**b. The Parahippocampal Place Area (PPA).** The PPA is defined functionally as the region adjacent to the collateral sulcus in parahippocampal cortex that responds significantly more strongly to images of scenes than objects (Epstein & Kanwisher, 1998). The PPA responds to a wide variety of scenes, including indoor and outdoor scenes, familiar and
unfamiliar scenes, and even abstract “scenes” made of Legos (Aguirre, Detre, Alsop & D'Esposito, 1996; Epstein, 2008; Epstein, 2005). The PPA is primarily responsive to the spatial layout of one’s surroundings: Its response is not reduced when all of the objects are removed from an indoor scene, leaving just the floor and walls (Epstein & Kanwisher, 1998). The PPA has also been shown to respond selectively to high-spatial frequency geometric shapes in humans and monkeys (Rajimehr et al. 2011), suggesting that the PPA may use such information for detecting scene details during place perception and navigation. By contrast, Bar and colleagues have proposed that the parahippocampal response to scenes does not reflect spatial layout, but rather the activation of a “context frame” representation that includes information about which objects typically appear in that context and where they are likely to be located relative to each other (Bar, 2004). However, the evidence for the context hypothesis is weak: the finding of greater response to strong-versus weak-context objects only replicates at slow presentation rates, is only reliable in a minority of subjects, and can be alternatively explained in terms of scene imagery (Epstein & Ward, 2010). On the other hand, consistent with the spatial layout hypothesis, studies using MVPA found that the PPA contains significantly more information about spatial layout; that is, whether a scene is “open” versus “closed” (Oliva & Torralba, 2001) than information about whether a scene is manmade (e.g., a city) or natural (e.g., a forest) (Kravitz, Peng & Baker, 2011; Park, Brady, Greene & Oliva, 2011). Evidence that the PPA is not only activated when information about spatial layout is processed, but that it is further necessary for this function, comes from patients with damage in or near the PPA, who have deficits in simple identification of scenes or landmarks (Aguirre & D'Esposito, 1999; Mendez & Cherrier, 2003), and difficulty more generally in knowing where they are (Epstein, De Yoe, Press & Kanwisher, 2001; Habib & Sirigu, 1987). This high response to spatial layout information is tantalizingly reminiscent of the “geometric module” (Cheng & Gallistel, 1984; Hermer & Spelke, 1996), inferred from behavioral data in which rats and human infants (and adults whose language system is tied up by a concurrent verbal task) rely exclusively on the layout of space, not on objects or landmarks, to reorient themselves in an environment once they are disoriented. However, we recently found that representations in the PPA are largely invariant to mirror-image reversals, a result that challenges its role in navigation and reorientation (Dilks, Julian, Kubilius, Spelke & Kanwisher, 2011). The precise role of the PPA in place perception and navigation is a topic of ongoing investigation (Oliva, this volume).

c. The Extrastriate Body Area (EBA). The EBA is a region on the lateral surface of the brain adjacent to (and sometimes partly overlapping with) visual motion area MT, which responds significantly more strongly to images of bodies than to images of objects or faces (Downing et al., 2001). This region responds equally to visually different images of bodies, from a photograph of a hand, to a photograph of a body (human or animal), to a line drawing and even a schematic stick figure of a person. The EBA is more involved in perceiving other people’s bodies, in a viewpoint-dependent manner (Taylor, Wiggett & Downing, 2010), than one’s own (Chan, Peelen & Downing, 2004; Saxe, Jamal & Powell, 2006), and is more engaged in the perception of the form/identity of bodies than in the actions they are carrying out (Downing & Peelen, 2011; Moro et al., 2008). Evidence that this region is not only activated during, but is also necessary for, the
perception of bodies comes from studies in which disruption of the EBA by a brain lesion (Moro et al., 2008) or by transcranial magnetic stimulation (TMS) (Pitcher, Charles, Devlin, Walsh & Duchaine, 2009; Urgesi, Berlucchi & Aglioti, 2004) impairs the perception of body form but not the perception of faces or of object shape (Pitcher et al., 2009). Recent evidence indicates that the EBA may consist of several subregions, and that these subregions of the EBA and neighboring cortex may respond differentially to distinct body parts (Bracci, Ietswaart, Peelen & Cavina-Pratesi, 2010; Op de Beeck, Brants, Baeck & Wagemans, 2010; Orlov, Makin & Zohary, 2010; Weiner & Grill-Spector, 2011). A recent review (Downing & Peelen, 2011) argues that the EBA, and another more ventral body-specific region, the fusiform body area, or FBA (Schwarzlose, Baker & Kanwisher, 2005), are not primarily engaged in a higher-level interpretation of the individual identity, emotional content, motion, or action goals, but rather extract a “cognitively unelaborated” visual representation of the shape and posture of the people in the current percept.

d. The Lateral Occipital Complex (LOC). In addition to the category-specific regions just described, a large region of lateral and inferior occipital cortex just anterior to retinotopic cortex (“LOC” for lateral occipital complex), and partially overlapping with some of the regions described above, responds more strongly to stimuli depicting shapes than stimuli with similar low-level features that do not depict shapes (Kanwisher et al., 1997; Malach et al., 1995). Common areas within LOC are activated by shapes defined by motion, texture, and luminance contours (Grill-Spector, Kushnir, Edelman, Itzchak & Malach, 1998), showing that representations of shape in LOC are quite abstract. Importantly, the LOC responds similarly to familiar and unfamiliar shapes (Kanwisher et al., 1997; Malach et al., 1995), suggesting that this region is not involved in matching to stored object representations, or semantic or verbal coding of the stimuli. Further support for the idea that LOC represents object shape, not semantic information about objects, comes from the fact that fMRI adaptation is not found in LOC across objects that are similar in meaning but differ in shape (Kim, Biederman, Lescroart & Hayworth, 2009). Several studies have implicated the LOC in visual object recognition by showing that activity in this region is correlated with success on a variety of object recognition tasks (Bar et al., 2001; Grill-Spector, Kushnir, Hendler & Malach, 2000), and indeed the location of LOC matches very nicely the location of the lesion in the famous ventral pathway agnostic patient DF (James, Culham, Humphrey, Milner & Goodale, 2003). Thus an investigation of the response properties of the LOC may provide important clues about the nature of the representations underlying object recognition.

MVPA and fMRI adaptation studies show that LOC contains fairly abstract representations of object shape. First, fMRI adaptation studies have shown that representations in the vicinity of LOC are partially invariant to changes in size and position, but largely specific to viewpoint and direction of illumination (Grill-Spector et al., 1999). Evidence on the nature of the shape representations in LOC comes from the fact that fMRI adaptation occurs in LOC between stimulus pairs that have different contours but the same perceived shape (because of changes in occlusion), but not between stimulus pairs with identical contours but different perceived shape (because of a figure-ground reversal) (Kourtzi & Kanwisher, 2001). Further, MVPA analyses show
that a posterior subregion of LOC (often called “LO”) contains representations that are more tied to the stimulus, whereas a more anterior subregion (called “pFs”) contains representations that are correlated with observer-specific perceptions of shape similarity (Haushofer, Livingstone & Kanwisher, 2008). Finally, another recent study found that the more posterior region LO showed sensitivity to mirror-image reversals of objects, while the more anterior region pFs did not, suggesting a hierarchy of object processing in which left–right information is represented at earlier (more posterior) stages in the hierarchy and invariance is then computed at later (more anterior) stages (Dilks et al., 2011).

In sum, the VVP contains a large multi-part region, the LOC, that responds strongly to object structure but that exhibits little selectivity for specific object categories, along with a small number of category-specific regions (for faces, places, and bodies). Efforts to date have not found regions of the VVP robustly selective for other categories (Downing, Chan, Peelan, Dodds & Kanwisher, 2006; Lashkari et al., 2011), with the exception of the VWFA (Baker et al., 2007; Cohen et al., 2000).

3. Specificity: Do category-selective regions contribute only to the perception of their preferred stimuli?

So far, we have argued that the FFA, PPA, and EBA are primarily if not exclusively engaged in processing their “preferred” stimuli (the stimulus class they respond most strongly to). However, each of these regions responds significantly (albeit weakly) to objects that are not in the preferred category (aka “nonpreferred” objects). Further, in what we consider the most important challenge to the claimed specificity of these regions, Haxby and colleagues (Haxby et al., 2001) reported that the spatial pattern of response across the FFA contains information about nonfaces, and that the pattern of response within the PPA contains information about nonscenes, and hence that “Regions such as the ‘PPA’ and ‘FFA’ are not dedicated to representing only spatial arrangements or human faces, but, rather, are part of a more extended representation for all objects.”

We too find that the FFA and PPA contain information about nonpreferred stimuli (Reddy & Kanwisher, 2007), and current physiological evidence (Tsao, Freiwald, Tootell & Livingstone, 2006) also indicates that face selective regions carry a small but significant amount of information about nonpreferred stimuli. However, the information that category-selective regions contain is much weaker for nonpreferred than preferred stimuli. As Haxby and colleagues (O’Toole, Jiang, Abdi & Haxby, 2005) noted, “preferred regions for faces and houses are not well suited to object classifications that do not involve faces and houses, respectively.” These findings raise two questions, which we address next.

First, is the information about nonpreferred objects in category-selective regions still present when subjects view cluttered displays more typical of real-world vision? Most analyses of the spatial pattern of the fMRI response within the VVP have been based on responses elicited by single cut-out objects on a blank background, presented at the
fovea. Of course real-world visual stimuli are not this simple: A typical visual scene contains multiple objects and complex background textures (i.e., “clutter”). A recent study (Reddy & Kanwisher, 2007) tested whether information about nonpreferred objects is present in a minimalist case of clutter, with two objects present simultaneously in the visual field (both on blank backgrounds). When single cut-out stimuli were shown one at a time, the pattern of response in the FFA contained considerable information about faces, and significant though weaker information about nonfaces. Similarly, the pattern of response in the PPA contained robust information about houses (which activate the PPA strongly, though not as strongly as a full scene) and significant but weak information about nonscenes. Crucially, however, when two objects were present at once, information about preferred stimuli was virtually undiminished from the single-object case, but information about nonpreferred stimuli dropped to insignificance (Reddy & Kanwisher, 2007). This study and later related studies suggest that category-selective regions may have little or no information about nonpreferred stimuli under natural (i.e., cluttered) viewing conditions.

Still, given that fMRI is bound to underestimate the information present in the full neural population code, it is possible that future physiological studies will reveal some information about nonpreferred stimuli in the FFA, the PPA, and similarly selective regions, even for the cluttered stimuli typical of real-world viewing. Thus the second and most important question is whether such information is used in the perception of those stimuli, or whether it is epiphenomenal (Williams, Berberovic & Mattingley, 2007). Some relevant evidence is available for the case of the FFA from the study of individuals with focal brain damage. Some of these individuals exhibit deficits only in face perception (i.e., prosopagnosia), with little or no deficit in object recognition, after damage to regions in or near the FFA, suggesting that even if the FFA contains information about nonfaces, this information is not necessary for object perception. Though no published case of acquired prosopagnosia has completely ruled out the existence of any other deficits beyond face perception (Garrido, Duchaine & Nakayama, 2008) using the most sensitive tests of object perception such as reaction time measures (Gauthier, Behrmann & Tarr, 1999), some cases come close (Sergent & Signoret, 1992; Wada & Yamamoto, 2001). However, because the locus and extent of lesions in humans is not under our control, an importantly complementary method for testing the functional specificity and causal role of cortical regions in perception is TMS. In TMS, a brief magnetic pulse is delivered to the scalp through a coil held next to the scalp, disrupting neural processing in the cortical region immediately beneath the coil. We can now precisely position the TMS coil to directly target specific cortical regions defined functionally within individual subjects. Although the FFA is too medial to be reached by TMS, another more lateral face-selective region (the OFA, discussed in more detail in Section 4) can be. Using this method, Pitcher and colleagues (Pitcher et al., 2009) showed that TMS to the EBA disrupted perception of bodies (Urgesi et al., 2004) but not faces or objects, TMS to the OFA disrupted perception of faces but not objects or bodies (but see Silvanto, Schwarzkopf, Gilaie-Dotan & Rees, 2010), and TMS to LO disrupted perception of objects but not bodies or faces. This striking triple dissociation suggests that category-selective regions play a causal role in the perception of their preferred stimulus class, but not their nonpreferred stimulus class. Thus, even if the pattern of
response across these regions contains some information about nonpreferred stimulus categories, the available evidence suggests that such information plays no detectable causal role in perception.

In sum, current evidence suggests that category-selective regions sometimes contain weak but significant information about nonpreferred stimuli, which is likely to be underestimated by fMRI. Nonetheless, results from neuropsychology and TMS are consistent with the hypothesis that any information about nonpreferred stimuli in category-selective regions is epiphenomenal (i.e., not causally involved in perception of those stimuli). It will be important in the future to test this hypothesis further with new data from patients, TMS, and other disruption methods, such as electrical microstimulation in macaque monkeys and humans (Afraz, Kiani & Esteky, 2006; Puce et al., 1999).

4. The function and structure of the whole ventral visual pathway

Of course no complex cognitive process is accomplished in a single brain region, and arguments for the specificity of the regions described above in no way preclude an important role for other brain regions. “Earlier” cortical regions such as primary visual cortex are obviously crucial in the perception of faces, places, and bodies, and “higher” areas (e.g., in parietal and frontal regions) are also probably necessary for information in the FFA, PPA, and EBA to be used by other cognitive systems and to reach awareness (Kanwisher, 2001). Further, none of these regions is the only one with its defining selectivity. Other category-selective regions have not been studied in the same detail as the FFA, PPA, and EBA, so their functions are less clear. However, the existence of multiple selective regions, and the growing evidence for a functional division of labor between them, raises the exciting possibility that we may ultimately understand how face recognition, for example, emerges from the joint activity of a number of functionally distinct regions. Next, we briefly review the literature on other functionally distinct regions engaged in face and scene perception.

For faces, selective responses are found not only in the FFA, but in many subjects also in a nearby but more posterior OFA (Gauthier et al., 2000), as well as other more anterior regions, such as the posterior superior temporal sulcus (pSTS) (Puce, Allison, Bentin, Gore & McCarthy, 1998), and anterior temporal pole (Rajimehr, Young & Tootell, 2009). Based on the more posterior location and generally lower selectivity of the OFA, it is often assumed to constitute an early stage of face perception (or face detection), which is then followed by continued processing in more anterior regions (e.g., the FFA and pSTS) (Haxby, Hoffman & Gobbini, 2000; but see Rossion, Hanseeuw & Dricot, 2012). Consistent with this picture, i) the right OFA has been more implicated in the representation of the parts of a face, including the eyes, nose, and mouth, than the configuration of those parts (Liu et al., 2010; Pitcher, Walsh, Yovel & Duchaine, 2007), whereas the FFA is sensitive to both face parts and their overall configuration in the face (Liu et al., 2010) and ii) the OFA is more sensitive to mirror-image reversals than is the FFA or pSTS (Axelrod & Yovel, 2012). By contrast, the face-selective region in the pSTS has been implicated in the representation of more dynamic high-level face and
social information, including eye, mouth and head movements (Carlin, Rowe, Kriegeskorte, Thompson & Calder, 2011; Fox, Moon, Iaria & Barton, 2009; Haxby et al., 2000; Pitcher, Dilks, Saxe, Triantafyllou & Kanwisher, 2011; Puce et al., 1998) and facial expression (Phillips & David, 1997; Winston, Henson, Fine-Goulden & Dolan, 2004). In one of the most striking functional dissociations within the face system, the face-selective region in the pSTS responds about three times as strongly to movies of faces (but not movies of bodies or objects) as to static snapshots taken from those face movies, whereas the FFA responds the same to movies and snapshots (Pitcher et al., 2011; see also Puce et al., 1998).

For scenes, selective responses are found not only in the PPA, but also in retrosplenial complex (RSC), and the transverse occipital sulcus (TOS). Like PPA, RSC is primarily responsive to the spatial layout of one’s surroundings (Dilks et al., 2011; Epstein, 2008; Kravitz et al., 2011; Park & Chun, 2009), with a recent study reporting only spatial layout information, not object information, in RSC (Harel, Kravitz & Baker, 2012). However, in contrast to the ongoing debate about the PPA’s precise role in place perception and navigation, most studies find clear evidence that RSC plays a role in navigation. For example, a fMRI adaptation study (Baumann & Mattingley, 2010) has shown that this region encodes heading direction. Further, RSC is sensitive to left-right information in scenes (i.e., mirror-image reversals of a scene), which is presumably important for navigation (Dilks et al., 2011). Finally, patients with RSC damage have been reported to recognize salient landmarks but not use these landmarks to orient themselves or to navigate through a larger environment (Takahashi, Kawamura, Shiota, Kasahata & Hirayama, 1997). TOS is the least studied scene-selective region, but preliminary data from our lab suggests it is causally involved in scene perception: TMS over TOS impaired discrimination of scenes but not faces (Julian, Kanwisher & Dilks, 2012).

The existence of multiple face-selective regions, and multiple scene-selective regions offers the exciting prospect of taking apart the process of face and scene perception by understanding the functional division of labor between the various regions within each system. An important part of this story concerns the connectivity of the different regions within each system, and between those regions and the rest of the brain. Evidence on this important question is sparse, however, because neither of the two methods currently available in humans can answer these questions definitively. Resting-state fMRI is intriguing, but can reveal strong correlations between regions known not to be directly connected (Tian et al., 2007); Diffusion tractography is subject to ambiguities both in tracing connectivity from specific functionally-identified grey matter regions into the underlying white matter, and in tracing specific connections through white matter (rather than simply following known major fiber bundles that run through the VVP, e.g., the ILF and IFOF). Indeed, preliminary evidence from these methods does not fully agree: While both diffusion tractography and resting-state fMRI agree that the nearby OFA and FFA are connected, connections between the FFA and the face-selective region in the pSTS have been found with resting-state fMRI (Turk-Browne, Norman-Haignere & McCarthy, 2010), but not with diffusion methods (Gschwind, Pourtois, Schwartz, Van De Ville & Vuilleumier, 2011). It remains to be determined whether methods for
discovering specific anatomical connections in humans will ultimately be able to discover the precise connectivity of specific subregions of the VVP.

Perhaps the biggest open question concerning the functional organization of the VVP is whether the functionally-distinctive regions identified here are best thought of as discrete processors, or whether it makes more sense to consider the broader region that contains them as a single processor, in which each of these regions simply constitutes a local peak in the functional response. On the latter view, the question would still remain of why that landscape would contain the particular replicable configuration it does across the VVP, and what if any are the dimensions represented by axes of this broader “map” (Op de Beeck, Haushofer & Kanwisher, 2008; Kanwisher & Schwarzlose, 2008). Some of the locations of particular regions in the VVP may be explained in terms of a center-periphery map (Hasson, Levy, Behrmann, Hendler & Malach, 2002), or a representation of real-world object size (Konkle, 2011). Further, widely noted and intriguing aspects of the structure of the VVP are that many of the object- and category-selective regions come in pairs, with one on the ventral surface and one on the lateral surface (Hasson, Harel, Levy & Malach, 2003; Schwarzlose et al., 2008), and that body-selective and face-selective regions tend to be close and sometimes overlapping with each other (Kanwisher & Schwarzlose, 2008; Weiner & Grill-Spector, 2011). One clue into the question of whether the ventral pathway is best thought of as a single representational space, or a set of at least partially distinct processors, may come from anatomy: To the extent that the different regions discussed above have distinctive connectivity and cytoarchitecture, that would support the interpretation of these regions as distinct entities. Indeed, recent evidence indicates that face selectivity within the fusiform gyrus can be predicted from connectivity with the rest of the brain (Saygin et al., 2011), and that at least one face-selective region in the fusiform gyrus may have a distinctive cytoarchitecture (Caspers et al., 2012).

5. Why have selective regions in the first place?

Perhaps the deepest question raised by the work on the VVP is this: Why do some visual categories get their own private piece of real estate in the VVP, while others apparently do not (Downing et al., 2006; Lashkari et al., 2011)? To think clearly about this question, we need to consider what computational advantages are afforded by functional specialization in the first place. To be detected by fMRI, functional specializations must have two properties: i) selectivity of the response of neurons to the relevant information (e.g., face selectivity), and ii) spatial clustering of selective neurons. These phenomena are related but distinct (Ohki, Chung, Ch'ng, Kara & Reid, 2005), and will be discussed in turn in this final, highly speculative section.

a. Selectivity/Sparseness. The advantages of selectivity, or “sparseness,” in neural coding have been widely noted (Barlow, 1995; Foldiak & Young, 1995; Olshausen & Field, 2004). If a given object is coded by the activity of a small subset of the available neurons, then interference is minimized in two important senses. First, it is possible to represent multiple objects simultaneously with minimal ambiguity, because the neural codes for different objects are unlikely to overlap. Thus, we can perceive a face and place
simultaneously without the two representations colliding (Reddy & Kanwisher, 2007). Perhaps this is one reason we have neural populations selectively responsive to faces, places, and bodies: to provide “private lines” of communication about particularly important classes of stimuli that are protected from crosstalk of other irrelevant information.

Second, the use of sparse codes can also reduce interference from learning, enabling us to learn new exemplars of one class of objects without altering stored information about another class of objects. With one neural population to represent faces and a nonoverlapping neural population to represent the spatial layout of places, we can learn new faces without disrupting our memories of places and vice versa. From this perspective, we may expect to find relatively sparse codes for classes of information characterized by continual lifelong learning (like faces and places).

Third, building specialized brain regions, and precise connectivity linking them to other brain regions, could bootstrap development by essentially hardwiring constraints on inductive inference. For example, if information in faces provides the key input required for learning about other people’s minds, then perhaps the most efficient way to construct the machinery for thinking about other minds is to hardwire a face area and connect it to another available region, which will then have the constrained input it needs to construct the circuits necessary for social cognition. Evidence against this particular hypothesis comes from the recent finding that congenitally blind individuals show the same location and pattern of activation as sighted subjects when thinking about other people’s thoughts, even though input from the FFA is likely very different or nonexistent in these people (Bedny, Pascual-Leone & Saxe, 2009). Nevertheless, the general idea that specialized brain regions and their connections may serve as constraints on development is worth considering in other cases.

A fourth possible advantage of relatively sparse codes is metabolic rather than computational: Less energy is required if fewer neurons are firing. From this perspective, the greatest lifelong energy savings would come about if sparse codes were available for classes of stimuli that occur most frequently (Foldiak & Young, 1995). Thus, even from a purely metabolic perspective, it makes sense to use relatively sparse codes for faces, places, and bodies, because they are among the most frequently encountered visual stimuli.

In sum, sparse codes, in which information is represented by a relatively small percentage of the available neurons, each with relatively high selectivity, have certain advantages. At the same time, sparse codes have well-known disadvantages, such as greater susceptibility to damage (because of the smaller number of neurons involved in any given representation), and a smaller number of possible patterns that can be held (one at a time) by a fixed number of neurons. The speculation here is that these disadvantages are outweighed by the particular advantages in the coding of biologically important stimuli like faces, places, and bodies: i) reduction of interference or crosstalk when multiple stimuli must be represented simultaneously, ii) the ability to learn new information about one stimulus class without disrupting stored information about another class, iii) bootstrapping the development of other regions, and iv) the potential energy efficiency of coding the most frequently-encountered stimuli through the activity of the smallest number of neurons.
b. Spatial Clustering. The second property implied by functionally selective regions detected by fMRI, after selectivity of neurons, is spatial clustering of those neurons. Spatial clustering of functional properties is a familiar phenomenon in the brain, found not only in retinotopic, somatotopic, tonotopic, and other-topic maps that follow the organization of the receptor surface, but also in the organization of functional information that is computed de novo, like orientation columns in primary visual cortex and chromotopic maps in posterior inferotemporal cortex (Conway & Tsao, 2009). Spatial organization is such a pervasive and familiar property of the cortex that we can easily forget to ask ourselves why it occurs. This mystery has been articulated most clearly (Chklovskii & Koulakov, 2004) as follows: “Imagine taking a cortical area containing a map and scrambling neurons in that area, while preserving all the connections between neurons. Because the circuit remains unchanged, the functional properties of the neurons remain intact. Then the scrambled region without a map is functionally identical to the original one with the map.” Given that the identical circuit can be constructed in a spatially clustered or spatially scrambled version, why does spatial clustering occur?

This question is sharpened by the facts that the strong spatial clustering seen in some systems, such as orientation-selective cells in cat visual cortex, is not found in other very similar systems, such as orientation-selective cells in rodent visual cortex (Ohki & Reid, 2007), and further by the fact that in the rodent olfactory processing pathway, the precise spatial clustering (and odorant specificity) constructed in the olfactory bulb is thrown away in the next stage of processing, the piriform cortex (Stettler & Axel, 2009).

Chkovskii and Koulakov (2004) argue that the need to minimize wiring length (for developmental, metabolic, and conduction delay reasons) must be a fundamental constraint in the nervous system that produces spatial clustering of neurons that are densely connected to each other. To the extent that this wiring-length minimization principle is an important determinant of cortical organization, it suggests that we may find functional specialization in focal cortical regions for functions that are implemented in circuits for which the neurons have to be densely connected to each other. A testable prediction of this idea is that neurons within face-selective patches of monkey cortex must be richly interconnected, either directly, or via webs of inhibitory interneurons found in those same regions. A further prediction of the axon-length minimization principle is that to the extent that readout of a neural code (by the next stage of processing) requires convergence of multiple inputs on a particular neuron, it may be easier to read out a population code represented in a focal region of cortex where those inputs can all conveniently converge on a common output neuron. In a different vein, the functional significance of spatial clustering in the cortex may derive from the requirement to selectively modulate a given functional circuit by way of nonsynaptic diffusible messenger molecules that can spread a few millimeters through the cortex.

c. Functionally-specific Cortical Regions for Computationally Different Problems? Of course, the classic argument for functional specialization is that efficiency can be gained by division of labor (Rueffler, Hermisson & Wagner, 2012) when the computational requirements differ across tasks (Marr, 1982). Indeed, especially for the case of faces and places, both theoretical considerations and extensive empirical evidence suggest that
different kinds of representations are extracted from these stimulus classes and different uses are made of the resulting information (Cheng & Gallistel, 1984; Hermer & Spelke, 1996; Mckone & Robbins, 2010). On the other hand, the observed clustered selectivity for faces, places, and bodies does not in itself imply qualitative differences in the computations and representations entailed in the perception of one of these categories versus another. We might have functional selectivity of the relevant neuronal responses for each category without any fundamental differences in the kinds of computations conducted for each, just as we see in retinotopic cortex, where completely nonoverlapping pools of neurons code for visual information in one visual field location versus another, but fundamentally similar computations are conducted by each. A crucial question for the enterprise of using functional specificity of the brain to infer fundamental components of the mind will therefore be: which cortical selectivities reflect fundamentally different underlying cognitive processes and which simply reflect convenient compartmentalization of similar processes?

6. Conclusions

Over the past fifteen years, fMRI studies have taught us a great deal about the functional organization of the VVP in humans. We have learned that the machinery that conducts visual object recognition in humans is not a homogeneous mass of tissue, but instead a richly structure system composed of functionally distinct regions, each found in approximately the same location in every healthy subject. Exciting directions for future research will exploit a suite of powerful new methods, including the ability to relate the “representational dissimilarity matrices” extracted in each region (or the whole ventral pathway) via fMRI with representational spaces derived from behavioral and monkey single-unit data (Kriegeskorte et al., 2008), and the ability to link functionally-defined regions and functional profiles with cytoarchitecture (Caspers et al., 2012), and connectivity (Saygin et al., 2011).

Yet many fundamental questions have proven to be difficult or impossible to answer with current methods available in humans. What information is represented in each region at the spatial and temporal resolution of actual neural responses (Freiwald & Tsao, 2010)? What neural circuits extract this information? What is the causal role of each region in perception? What is the connectivity among these various regions, and between each of them and the rest of the brain (Moeller, Freiwald & Tsao, 2008)? Further, how do these regions get wired up in the brain in development, and what are the relative roles of experience (Baker et al., 2007; Srihasam, Mandeville, Morocz, Sullivan & Livingstone, 2012), and genes (Duchaine, Germine & Nakayama, 2007; Sugita, 2008; Turati, Bulf & Simion, 2008; Zhu et al., 2010) in this process? One of the most exciting developments of the last decade has been the discovery of face-selective (Tsao et al., 2006) and place-selective (Nasr et al., 2011) regions in the temporal lobes of macaques. These discoveries offer the possibility that the questions that have proven most intractable in humans will be answered by work on macaques, where it is possible to provide much richer characterizations of neural representations and their underlying circuits, the causal roles of these circuits in behavior, the structural correlates of functionally defined
regions, the interplay of genes and experience that wire these regions up during development, and the interactions among those regions during task performance.
Figure Legends

Figure 1. Inflated brain from three representative subjects showing regions specifically involved in the perception of faces (blues), places (pinks), and bodies (green). Dark blue = FFA; Purplish blue = pSTS; Light blue = OFA. Magenta = PPA; Light purple/pink = RSC; Reddish pink = TOS; Green = EBA. Each of these regions can be found in a short functional scan in essentially every healthy subject. LOC is not shown here; It is a very large region generally responsive to any object shape, and hence both of its subregions (LO and pFs) overlap partially with some of the regions shown here. The VWFA (left hemisphere) and FBA (partially overlapping with FFA) are also not pictured.
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